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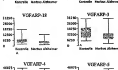
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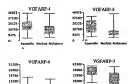
[Fortsetzung auf der nächsten Seite]

(54) Title: METHOD FOR DETECTING CHRONIC DEMENTIA DISEASES, AND CORRESPONDING PEPTIDES AND DE-TECTION REAGENTS

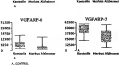
(54) Bezelchnung: VERFAHREN ZUM NACHWEIS CHRONISCH-DEMENZIELLER ERKRANKUNGEN, ZUGEHÖRIGE PEPTIDE UND NACHWEISREAGENZIEN







WO 02/082075



(57) Abstract: The invention relates to defined peptides and the quantitative determination thereof in biological samples from patients suffering from Alzheimer's disease, in relation to the concentration thereof in a control group. The invention also relates to the use of said peptides for therapeutic purposes. The inventive peptides come from a protein precursor having the corresponding gene and are processed in a specific manner and modified in a post-translational manner. Changes in the concentrations of said peptides indicate Alzheimer's disease, and the direction of the change in concentration is specific for each peptide. Alzheimer's disease is detected by identifying the peptides individually or in groups. The invention can also be used to control the course of Alzheimer's disease, for the prognosis thereof and for the development of therapeutic agents to combat the same.

(57) Zusammenfassung: Die vorliegende Erfindung betrifft definierte Peptide und deren quantitative Bestimmung in biologischen Proben von Patienten, die an Morbus Alzheimer erkrankt sind, relativ zu deren Konzentration in einer Kontrollgruppe. Ausserdem betrifft die Erfindung die Verwendung der Peptide zu therapeutischen Zwecken. Die erfindungsgemässen Peptide entstammen aus einem Proteinvorläufer mit dem korrespondierenden Gen und sind in spezifischer Art und Weise prozessiert und postranslational modifiziert. Änderungen der Konzentrationen dieser Peptide zeigen eine Morbus Alzheimer Erkrankung an. Dabei ist die Richtung der Konzentrationsänderung für jedes dieser Peptide spezifisch. Der Nachweis von Morbus Alzheimer erfolgt durch eine Identifizierung der Peptide einzeln oder in Kombinationen. Die Erfindung findet darüber hinaus Verwendung zur Verlaufskontrolle von Morbus Alzheimer, zur Prognose und zur Entwicklung von Therapeutika gegen Morbus Alzheimer.





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(54) METHOD FOR DETECTING CHRONIC DEMENTIA DISEASES, AND CORRESPONDING VGF PEPTIDES AND

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ABSTRACT

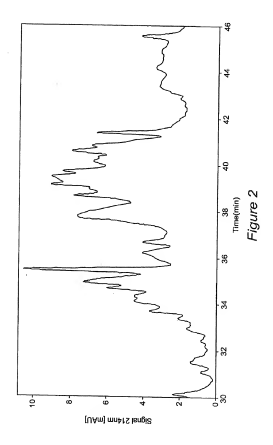
The invention relates to defined peptides and the quantitative determination thereof in biological samples from patient's suffering from Alzheimer's disease, in relation to the concentration thereof in a control group. The invention also relates to the use of said peptides for therapeutic purposes. The inventive peptides come from a protein precursor having the corresponding gene and are processed in a specific manner and modified in a post-translation manner. Changes in the concentrations of said peptides indicate Alzheimer's disease, and the direction of the change in concentration is specific for each peptide. Alzheimer's disease is detected by identifying the peptides individually or in groups. The invention can also be used to control the course of Alzheimer's disease, for the prognosis thereof and for the development of therapeutic agents to combat the same.

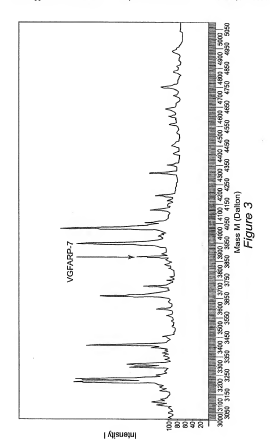
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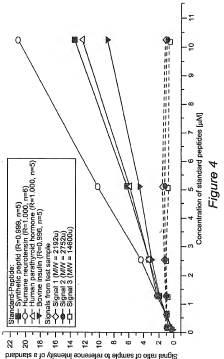
Y12661

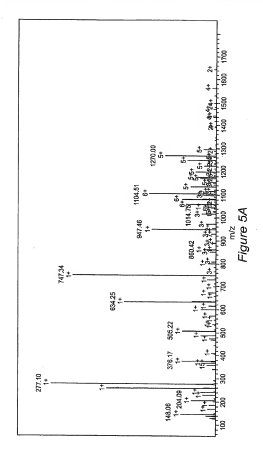
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VGFARP-2	APPGRPEAQPPPLSSEHKEPVAGDAVPGPKDGSAPEVRGA.
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VGFARP-5	GRPEAQPPPLSSEHKEPVAGDAVPGPKDGSAPEV
VGFARP-6	GRPEAQPPPLSSEHKEPVAGDAVPGPKDGS
VGFARP-7	GRPEAQPPPLSSEHKEPVAGDAVPGPKGA.
VGFARP-19	GRPEAQPPPLSSEHKEPVAGDAVPGPKDGSAPE
VGFARP-20	GRPEAQPPPLSSEHKEPVAGDAVPGPKDGSAP.
VGFARP-21	GRPEAQPPPLSSEHKEPVAGDAVPGPKDGSAPEVRGARM
VGFARP-10	PGFKDGSAPEVRGA
VGFARP-22	TDRPASPASPABEL
VGFARP-15	.rl-GPKDGSAP-r2
VGFARP-23	r7-HKEPVAGD-r8
VGFARP-24	r9-APSGSQQG-r10
Y12661 1	11261 117 ETVROCISIERAGEPERARPRETTERNOERASDESELEALASILQEIROSSESSAKROCETAABETETRIFILITRVILESPGERAVRASWGEPOARVPBRAEJFPRARSGFO NA 103778 117
VGFARP-25	SQTHSLPA <u>RES</u> BPEPARPPTFPTPENGPERSDPSEEL
VGFARP-26	QELROFSPSSA
VGFARP-27	TIT-FRANKERIA
Y12661 2 NM_003378 2	11266 233 ARMPOGEPETHKFGEGVISPKTHLGENAPLERARNOGVAAPPRARARSBALLGGSEAGERLLGGGLAQVEAGRGGEATRQAAAGERLADLAGDLILOTILGGGRAGGEG RM. 003378 233 **********************************

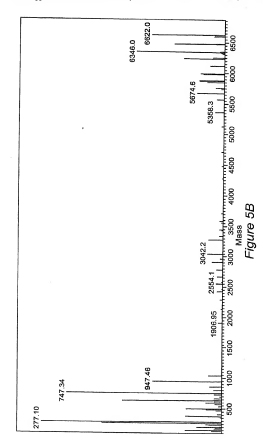
349 RGIQEAAREREBEAAQERRYGGERYGGERYGGERYGERAAAAAAAAAAAAAAAAAA	. LOEAABERRESAREEEBARGERRGGERVGEEDEAABAAEAEABEADEAGRARQNALLFABEBEDGEAGAED.	GLQBABERRSAREEEBA	GLQEAAEBRESARESEBARQE	. GGEERVGEEDEEAARA-FAEARBRARQNALLFAEEEDGEAGABED	GGEERVGEEDEEAARP-FAEARFRONALI.	GEERVGEEDBEAAEAAEAAEAAEAAEAAEAAEAAEAAEAAEAAEAAEAA	SQEETPGHRAKEAEGTEEGGEEEDDEEMDPQTIDSL	SQEETPGHRKEAEGTEEGDBEMDPQTIDSLIELSTKIH—>	SQEETPGHRRKEABGTEGGEEGDD5EMDPQTIBSTKLH→	↑ HTXLS	r13-AEERESAR-r14	r3-EDEEAABA-r4	.r5-EBMDPQTI-r6.	11261 465 IRADOWSIIENVERGARKKARARPENDARAPHWASPOPPERPROARENTERPROARENTERPROTOPPERPROARENTHALLPSSARFORM AND 461 TO THE TOTAL OF THE TOTAL	172661 580 BAQARHAQDERARAERALOROGELEWYIEHULARAE
FIGLOEARERESA 9 ************	LQEAAEERESA	GLQEAAEERESA	.GLQEAAEERESAI								r13-AEERESA			LPADDVVSIIEBV ************************************	EAQARHAQQEEAE
K12661 349 WM_003378 349	GFARP-11	GFARP-28	GFARP-29	GFARP-30	'GFARP-31	GEARP-32	GEARP-33	GEARP-12	GEARP-13	'GEARP-34	GEARP-35	GEARP-16	VGFARP-17	112661 466 NM 003378 466 VGFARP-36 VGFARP-37 VGFARP-12 — VGFARP-34 — VGFARP-38	X12661 580

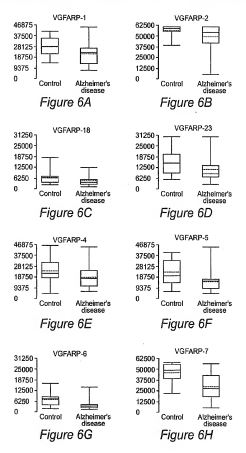


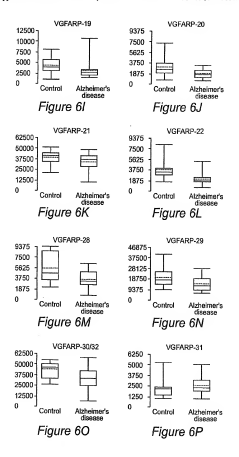


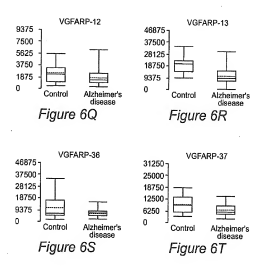












METHOD FOR DETECTING CHRONIC DEMENTIA DISEASES, AND CORRESPONDING VGF PEPTIDES AND DETECTION REAGENTS

[0001] This is a continuation-in-part (CIP) application of International Application PCT/DE02/01376 with an international filing date of Apr. 8, 2002, now abandoned.

FIELD OF THE INVENTION

[0002] The invention relates to a method for detecting a chronic dementia disease or a predisposition to a chronic dementia disease, in particular Alzheimer's disease or related neurological diseases, e.g. Lewy body dementia or vascular dementia. The invention further relates to peptides which have been found for detecting the presence of these diseases, for monitoring the course of the diseases and of the grade of the diseases. In addition, the invention relates to detection reagents such as antibodies and nucleic acids and the like, via which these peptides or the corresponding nucleic acids can be detected. The invention further relates to pharmaceutical applications which comprise VGF, VGF peptides, VGF antibodies, VGF nucleic acids, VGF protein antagonists, VGF protein agonists, VGF peptide agonists or VGF peptide antagonists for the therapy or prophylaxis of neurological diseases, especially of Alzheimer's disease. The invention further relates to methods for identifying patients with neurological diseases, especially Alzheimer's disease, who are suitable for taking part in clinical studies to investigate these diseases.

[0003] The peptides comprise fragments of the VGF protein, which is also called neuroendocrine specific protein VGF. The abbreviation VGF is also used in the literature for the protein "vaccinia growth factor" or for "vaccinia virus growth factor" and for "vascular permeability factor", these proteins not corresponding to the VGF protein to which the invention relates.

BACKGROUND OF THE INVENTION

[0004] Demontia diseases represent an increasing problem industrialized countries because of the higher average life expectancy. Dementia diseases are in most cases incurable and make long-term care of the patients necessary. About half of these patients receive impatient care. More than 60 dementia diseases are known, including diseases associated with manifestations of dementia.

[9065] However, Alzheimer's disease (AD) accounts for about 65% of these, and the diagnosis and therapy thereof is therefore of great importance. Besides Alzheimer's disease, the following non-Alzheimer's dementias are known, inter aliz vascular dementia, Lewy body dementia, Biaswanger dementia, and dementia diseases which occur as concomitant effects of other disorders such as Parkinson's disease, Huntington's disease, Pick's disease, Gerstmann-Sträussler-Scheinger disease, Kruzzfeld-Jadob disease etc.

[906] Alzheimer's disease is a neurodegenerative disease distinguisted by the following symptoms decline in distinguisted by the following symptoms decline in learnal abilities, confusion and diminished ability to look in a fart themselves. A greatly restricted short-term memory particular is characteristic of Alzheimer's disease, whereas each other confusion of the distantage of Alzheimer's disease, whereas the patient's memories of the distantage, e.g. of histography children of the distantage of the primary and applications of the distantage in the primary amanifested intent and morphological changes in the brain amanifested intent and morphological changes in the brain amanifested intent and

the farm of amyloid deponists and degenerated nerve cells. The morphological changes can be diagnosed histologically after the patient's death and are as yet the only reliable detection of the disease. These histopathological diagnoses are based on criteria fixed by the Consortium to Establish a Registry for Athelimer's Disease (CERAD). The following criteria-based diagnostic systems are currently used to diagnose Athelimer's diseases: the International classification of Diseases, 10th revision (ICD-10), the Diagnostic and Statistical Manual of Mental Disorders, 4th edition (DSM-1V) of the American Psychiatric Association, and the Work Group criest drawn up by the National Institute of Work Group criest drawn up by the National Institute of National Statistics of National Communicative Disorders Association NINOUS-ARDEN (NINOUS-ARDEN).

[0007] These systems use a number of neuropsychological tests in order to diagnose Alzheimer's disease, but not objectively measurable clinical parameters.

[0008] Diagnosis of Alzheimer's disease is also difficult because it, just like the other dementia diseases, has an insidious onset and is associated with slowly progressive destruction of nerve cells in the brain.

[0009] At present, no causal therapy is available for the textament of Abheimer's disease. The disease is merby treated symptomatically, e.g. by administration of neutoriasmities used as a scytyleholine. Further possible therapeatic strategies being tested at present are the administration of antioxidatas, of radical sexwangers, of calinochannel blockers, of antiinflammatory substances, of secretace inhibitors, of anti-myloid antibodies etc., and immunization against amyloid peptides. However, no causal therapy of this disease is yet possible.

SUMMARY OF THE INVENTION

[0010] The invention is based on the object of avoiding the prior at disadvotages in the diagnosis of Abzheimer's disease and of providing a method which can be used early and reliably for detecting chronic demonth afterses, especially Abzheimer's disease. It is additionally based on the object of providing a movel therapy for the treatment of Abzheimer's disease because, at present, only unsatisfactory therapeutic approaches to the treatment of Abzheimer's diseases are available.

DESCRIPTION OF THE DRAWINGS

[0011] FIG. 1: Alignment of the VGFARP peptides with the two known VGF proteins, corresponding to the database accession No. NM_003378 and Y12651, e.g. Seq. IDs 43 and 44

[0012] FIG. 2: Reverse phase chromatography for separation and enrichment of VGFARP peptides from cerebrospinal fluid

[0013] FIG. 3: Mass spectrometric measurement (MALDI) on VGFARP-7 (SEQ ID NO:7) as example

[0014] FIG. 4: MALDI as relatively quantifying mass spectroscopic method

[0015] FIG. 5: MS/MS fragment spectrum of the peptide VGFARP-13 (SEQ ID NO:11) as example

[0016] FIGS. 6a:-C: Box-whisker plots for quantitative comparison of the concentrations of VGFARP-1(SEO ID

NO:1), VGFARP-2(SEO DI NO:2), VGFARP-18(SEO DI NO:3), VGFARP-2(SEO DI NO:3), VGFARP-4(SEO DI NO:4), VGFARP-2(SEO DI NO:2), VGFARP-2(SEO DI NO:2), VGFARP-2(SEO DI NO:2), VGFARP-3(SEO DI NO:2), VGFARP-3(SEO DI NO:2), VGFARP-3(SEO DI NO:2), VGFARP-3(SEO DI NO:4), VGFARP-4(SEO D

DETAILED DESCRIPTION OF THE INVENTION

[0017] Definitions:

[0018] VGF Proteins or Peptides (SEQ ID NOS:44 and 43) Corresponding to Accession Nos. NM-003378 and Y12661: (SEQ ID NOS:46 and 45, Respectively)

[0019] The peptides (SEQ 1D NOS:43 and 44) derived from the nucleic acid sequences NM-003378 and Y12661 (SEQ ID NOS:44 and 43, respectively) are also referred to as VGF proteins and include all naturally occurring alleles, mutants and polymorphisms of VGF proteins, and tissuespecifically expressed VGF variants. Included in particular are also the VGF variants which occur because of diseases or as a result of neurological diseases, especially chronic dementia diseases, especially Alzheimer's disease. There is inclusion both of VGF proteins with and without signal sequence, proforms of VGF proteins which have not yet been processed, and already processed VGF proteins, soluble VGF proteins and membrane-associated VGF proteins, where the membrane-associated VGF proteins may be linked both via transmembrane amino acid sequences to a cell membrane or organelle membrane and via a posttranslational modification, e.g. a glycosyl-phosphatidylinositol (GPI) anchor. Also included are variations of the VGF sequence which [lacuna] by alternative splicing, by alternative translation starting and termination points, by RNA editing, by alternative post-translational modifications, and other VGF protein variants arising through naturally occurring mechanisms.

[0020] VGFARP Peptides:

[0021] VGF peptides and VGF peptide variants are referred to hereinafter as VGFARP (VGF Alzheimer related peptide) peptides. VGFARP peptides may be derived from both the VGF sequences mentioned at the outset (NM_ 003378=Seq. ID 43 for the protein and Seq. ID 45 for the DNA) and Y1266-Seq. ID 44 for the protein and Seq. ID 46 for the DNA) and from other VGF protein variants possibly occurring in nature. In addition, VGFARP peptides may include two point-mutated, two deleted and/or two additionally internally inserted amino acids, and/or N-terminal and/ or C-terminal extensions. However, in these cases they must retain at least 8 amino acids from the VGF protein sequence. VGFARP-39 (SEQ ID NO:36) is an exception from this rule, as VGFARP-39 (SEQ ID NO:36) has only a legth of 6 amino acids. The only amino acids suitable as N- or C-terminal extensions are those occurring in the VGF protein sequence at this sequence position in the VGF proteinpequites derived from naturally occurring VGF proteinphistos and from naturally occurring VGF mutants are also referred to as VGFARP peptides. VGFARP peptides may also exist with post-translational modifications such as, for example, glycosylations and phosphorylations and/or in chemically modified from, preferably as peptide oxides. For example, VGFARP-12 (SGD ID NO:10) has been identified both as non-oxidized and as oxidized peptide.

[9022] Chemically or Post-Translationally Modified Peptides;

[0023] A chemically or post-translationally modified peptide may consist both of D- and of L-amino acids, and of combinations of D- and L-amino acids. These pentides may additionally comprise unusual amino acids, i.e. amino acids which do not belong to the 20 standard amino acids. Examples of unusual amino acids are, inter alia: alphaaminobutyric acid, beta-aminobutyric acid, beta-alanine, beta-aminoisobutyric acid, norvaline, homoserine, norleucine, gamma-aminobutyric acid, thioproline, 4-hydroxyproline, alpha-aminoadipic acid, diaminobutyric acid, 4-aminobenzoic acid, homocysteine, alpha-aminopenicillanic acid, histamine, ornithine, glycine-proline dipeptide, hydroxylysine, proline-hydroxyproline dipeptide, cystathionine, ethionine, seleno-cysteine. Possible post-translational or chemical modifications are, inter alia, modifications of amino acid sequences by the following structures: linkage of free cysteine to a cysteine in the peptide sequence, methyl, acetyl, farnesyl, biotinyl, stearoyl, palmityl, lipoyl, C-mannosyl, phosphorus and sulfate groups, glycosylations, amidations, deamidations, pyroglutamic acid, citrulline etc.

[0024] Nucleic Acids:

[9025] Nucleis acids are regarded as being DNA, RNA, and DNA-RNA hybrid molecules both of natural origin and prepared synthetically or by recombination. Also included are chemically modified nucleic acids whole comprise modified nucleic acids whole comprise modified nucleic acids whole comprise ordified nucleic acids swing being in vivo stability, such as, for example, phosphoroutionates. Such a shallized nucleic acids are already used in the application of ribozyme, antisense and triplex nucleic acid etchniques.

[0026] Significance:

[0027] The term significant is used in the sense in which the term significance is used in statistics. In this patent application, an error probability of less than 90%, preferably 95% further preferably 95% is defined as significant.

[0028] Sensitivity:

[0029] Sensitivity is defined as the proportion of patients with the disease who acquire a positive diagnostic result in a diagnosis for the disease, i.e. the diagnosis correctly indicates the disease.

[0030] Specificity:

[0031] The specificity is defined as the proportion of healthy patients who acquire a negative diagnostic result in a diagnosis for the disease, i.e. the diagnosis correctly indicates that no disease is present.

[0032] It has surprisingly been found that only in samples of body fluids from patients suffering from Alzheimer's disease, especially in the cerebrospinal fluid, is the concentration of certain pertities chanced greatly relative to their concentration in control samples, and thus makes detection of Abzheimer's disease possible. Changes in the concentration of these peptides relative to their concentration in control groups indicate the presence of Abzheimer's disease and are therefore suitable for detecting lish disease with high sensitivity and specificity. Modulation of the VGF protein or VGFARP peptide concentration with the aim of adjusting the patient to normal VGF or VGFARP levels can thus be used therapeutically.

[0033] To achieve the object, the invention includes a method for detection of a neurological, in particular of a chronic dementia disease, in particular of Alzheimer's disease, or of a predisposition to such a disease by identifying one or more VGF peptides which are derived from the sequence having the Gene Bank accession No. NM_003378 or the accession No. Y12661 of the DNA Data Bank of Japan (Seq. ID 43 or 44), in a biological sample from an individual. Since these VGF peptides are presumably causally connected with the disease, the present invention also includes the use of these peptides for the therapy of Alzheimer's disease or related neurological diseases. These pentides or peptide fragments are referred to as VGF derived Alzheimer related peptides (VGFARP). The two VGF protein variants NM_003378 and Y12661 (SEQ ID NOS:44 and 43, respectively) differ only at 13 positions of their amino acid sequence and VGF peptides which make it possible to distinguish between Alzheimer's disease and the control group have been identified from both VGF proteins. The VGFARP peptides VGFARP-11 (SEQ ID NO:9), 32 (SEQ ID NO:29) and -44 (SEQ ID NO:41) are derived from the VGF variant with the accession No. Y12661(SEQ ID NO:43), and the VGFARP peptides VGFARP-25(SEQ ID NO:22), -30(SEQ ID NO:27), -31(SEQ ID NO:28), -36(SEO ID NO:33) and -37(SEO ID NO:34) are derived from the VGF variant with the accession No. NM-_ 003378(SEQ 1D NO:44). All the other VGFARP peptides can be derived on the basis of their amino acid sequence from both of the two VGF variants, Since VGFARP neptides derived from two different variants have already been identified, it must be assumed that further VGFARP peptides derived from these or other VGF variants also exist. The invention likewise relates to these VGFARP peptides.

[0034] To achieve the object, the invention indicates a method for the detection of Alzheimer's disease by determination of the relative concentration of at least one marker peptide in a biological sample from a patient compared with the concentration of the marker peptide in a control sample, in which the following points must be satisfied: 1. At least one VGFARP peptide or a peptide that is derived from the nucleic acids with the accession Nos. NM_003378 or Y12661 (Seq. 1Ds 45 and 46) or homologous sequences is used as marker peptide. 2. An increase or decrease specific for the particular marker peptide occurs in the concentration of the marker peptide in the patient's sample relative to the concentration of the marker peptide in the control sample. 3. A significant change in the concentration of the marker peptide in the aforementioned manner is regarded as a positive detection result for a neurological disease, preferahly Alzheimer's disease.

[0035] In this connection, it is possible in principle for a particular VGFARP peptide either to undergo only an increase in the peptide concentration in Alzheimer's disease patients. or it is possible in principle for this VGFARP peptide to undergo only a reduction in the peptide concentration of Alzheimer's disease patients. For a defined VGFARP peptide it is not possible for the VGFARP peptide of the undergood perturbation of Alzheimer's disease patient and to be reduced, relative to the control group, in another Alzheimer's disease patient and to be reduced, relative to the control group, in another Alzheimer's disease patient. As with virtually all medical diagnoses of disease patient. As with virtually all medical diagnoses of disease patient as with virtually all medical diagnoses of disease patient as with virtually all medical diagnoses of disease patient as with virtually all medical diagnoses of disease patients are possible in principle, i.e. that in a few individual cases an incorrect diagnosis takes place because the uncententation of the VGFARP peptides in Alzheimer's disease patients does not differ with bundred precent probability from the concentration of the VGFARP peptides in control samples. This problem can, however, be eliminated by mutiple controls.

[9036] Peptides which can be regarded as fragments of the VGF sequence are referred to as VGFARP peptides for the purposes of this invention. They include homologous reptides derived from VGF They include derivatives of manrally occurring alleles of these peptides and homologous mutants, especially point-mutated mutants with perforably on tone than two amino acids differing from VGF Preferred markers according to the invention are indicated in the sequence listing and thus named from VGFARP-1 (SEQ ID NC)1 to -7(SEQ ID NC). VGFARP-10 (SEQ ID NC)2 to -45(SEQ ID NC)2, corresponding to Seq. ID 1 to 42. The sequences of the VGFARP peptides are depicted in FIG. 1 and in Table 1. The assignment of the VGFARPspides to their respective Seq. ID No. is shown in Table 1.

[0837] The method of the invention comprises a method in which there is measurement of specific bounaries whose concentration is changed in neurodegenerative diseases, sepsentally in Adabetimer's disease, and which indicate the risk of the disease at n early class. This important in order to provide a reliable clinical marker for diagnosing these diseases.

[0038] It is possible and preferable for the concentration of VGFARP peptides in the sample, but also the characteristic pattern of occurrence of the plurality of particular VGFARP peptides, to be correlated with the severity of the disorder. These novel markets therefore make it possible to develop and monitor therapies for the treatment of Alzbeimero disease, because the course and any successful cure resulting from a therapy or a diministhed progression of the disease and be satisfasted. Effective therapy of Alzbeimer's disease is not possible at present, underlining the urgency for the provision of a railable detection method for Alzbeimer's disease, because reliable detection of the disease is a pre-condition for the development of a therapy.

[0039] Detection of VGFARP peptides additionally makes it possible in the framework of clinical studies to deep powel therapies for the treatment of Abzheimer's disease with high specificity to select only those patients suffering from Abzheimer's disease and not from other diseases. This is important for obtaining valid study results. Patients incorncetly diagnosed as Abzheimer's disease patients have a negative influence on the quality of the results of a study of Abzheimer's disease therapy. In addition, detection of VGFARP peptides makes it possible to stratify patients, i.e. the specific selection of subgroups of Abzheimer's disease patients who are especially satishly for particular Alzheimer's disease therapute strategies or clinical studies. [0040] There are marked changes in the concentrations of VGFARP peptides in Alzheimer's disease patients relative to healthy people. A further aspect of the invention is therefore a bringing of the VGFARP concentrations in Alzheimer's disease patients to normal concentrations. This method can be employed for the therapy of Alzheimer's disease or related neurological diseases. If the VGF protein or VGFARP peptide concentrations are elevated, the concentrations of these substances can be reduced by therapeutic administration of, for example, VGF protein- or VGFARP pentide-specific antibodies or VGF-specific antisense nucleic acids, ribozymes or triplex nucleic acids for VGFARP peptide antagonists, VGF protein antagonists. Substances which suppress the endogenous expression of VGF protein or the processing of VGF protein to VGFARP peptides can also be administered for the therapy. If the disease is caused by a deficiency of VGF protein or VGFARP peptides, therapeutic doses of VGF protein, VGFARP peptides, VGFARP peptide agonists or VGF protein agonists can be given. Endogenous production of VGF protein or VGFARP peptides can be increased by therapeutic administration of substances such as, for example, NGF, BNDF or NT-3 or other suitable substances, because these substances increase VGF expression. Substances which promote the processing of VGF protein to VGFARP peptides such as, for example, prohormone convertases such as, for example, PC1, PC2 or PC3, can also be employed therapeutically. Combination of different theraneutic strategies is. of course, also possible and sensible in some circumstances.

[0041] The invention therefore also encompasses the use of VGF proteins, VGFARP peptides, VGFARP peptide agonists and antagonists, VGF protein agonists and antagonists, anti-VGF protein antibodies, anti-VGFARP peptide antihodies, NGF, BNDF, NT-3, anti-NGF antibodies, anti-BNDF antibodies, anti-NT-3 antibodies and antibodies against receptors of said proteins for the direct or indirect modulation of the concentration of the VGF proteins and VGFARP peptides for the treatment of neurological diseases, especially Alzheimer's disease. Alternative to antibodies, it is also possible to use antibody fragments, antihody fusion proteins, or other substances which bind selectively to VGF proteins, VGFARP peptides, NGF, BNDF or NT-3. It is also possible as alternative to said proteins and peptides for fusion proteins of said proteins to be used. The invention further encompasses also the use of antisense nucleic acids, triplex nucleic acids and ribozymes which modulate the expression of said proteins and pentides. The invention additionally encompasses agonists and antagonists which modulate the activity of said proteins.

[0042] A further embodiment of the invention is the phameactual of manulation or chemical modification of the described peptides and mucleic acids to make it possible for them to cross the blood-brain barrier and/or the blood-CSF barrier more efficiently. They are thus made particularly suitable for therapeutic use. In order to achieve this, it is possible for example for VGF peptides, VGF proteins, uncleic acids, agoints or antagonists to be modified so that for example they become more lipophilic, favoring entry into the substanchoid space. This can be achieved by introducing hydrophobic molecular constituents or cles by "packaging" the substances in hydrophobic agents, e.g. liposomes. It is additionally possible for example for peptide sequences to be attached to these peptides, proteins, nucleic acids, agonists or antagonists, which favor crossing into the subarachnoid space or, conversely, impede crossing out of the subarachnoid space.

[9043] The invention also encompasses the administration of said therapeutic agents by various routes such a for example, as intravenous injection, as substance which can be administered orliky, as inhaltable gas or aerosol, or entrainistration in the form of direct injection into the sub-rachnoid space, or into tissue such as muscle, fal, brain ect. It is smoothly in the property of these therapeutic agents. For example, peptides or proteins administered orally can be protected by acid-resistant capsules from proteolytic degradation in the stomach. Very hydropholic substances can become more hydrophile and thus better suited for, for example, intravenous injections by suitable phramacutical processing etc.

[0044] A further embodiment of the invention is the use of VGEARP peptides or of VGF proteins for identifying receptors which selectively bind these molecules. These receptors can also be modulated by administration of agonisis or aniagonists, which is expedient for the therapy of neurological diseases, especially of Abzheimer's disease.

[0045] Owing to the large number of VGF peptides newly identified within the framework of this invention, it is possible for the first time to detect experimentally positions in the VGF protein at which processing of the VGF protein takes place in vivo. These processing sites comprise, hased on the VGF protein sequence of NM_003378 (SEQ ID NO:44), the following sequence positions: 371/372, 418/ 419, 479/480, 480/481, 481/482, 482/483 and 483/484. Based on the VGF protein sequence of Y12661(SEQ ID NO:43), the processing sites are as follows: 371/372, 419/ 420, 480/481, 483/484, 484/485 and 485/486. All experimentally identified processing positions represent dibasic positions, i.e. directly consecutive amino acids having positively charged amino acid side chains (arginine=R, lysine= K). Such sequence motifs are recognized and cut for example by prohormone convertases, with additional endoproteolytic deletion of the two basic amino acids. As the name of the prohormone convertases indicates, prohormones are converted by prohormone convertases to hormones, resulting in new bioactive substances (peptide hormones). Examples of biological active peptides which are generated in this way from their proforms are proNGF/NGF. pro BDNF/BNDF etc. [1]. Consequently, the VGFARP peptides of the invention represent peptide hormones which are suitable in connection with neurological diseases, preferably Alzheimer's disease, as points of attack for therapeutic agents. Modulation of the VGFARP peptide concentrations can thus he used for the therapy of neurological diseases, preferably Alzheimer's disease.

[0046] VGF Biology

[9047] The VGF proteins (VGF peptide precursor modcules) identified within the framework of this invention are synthesized as proteins about 68 kDs in size selectively in neuroendocrine and pursonal cells, with expression modern decreasing with increasing age [2]. Investigation of VGF gene-deficient mice revealed that important function in energy metabolism are affected [3]. VGF gene-deficient mice have a small body size, are hypermetabolic and hyperactive. VGF is also synthesized in the insulfin-producing islet cells of the pancerases. [0048] VGF was discovered on investigation of a rap phecohomocytom cell line (PGL 2cell line), and stimulo of this cell line with "enew growth factor" (NGF) brings about a 12- to 14-fold increase in the concentration of VGF [4, 5]. NGF is an important growth factor which regulates the differentiation of the peripheral and central nervous system. Further factors which regulate VGF expression are brain-derived neurotrophic factor (BDNF) and neurotropina. N(TA-3) [6]. VGF mRNA is regulated in volve perioral activity, neuronal injuries and by the biological rivthm (circatilla clock) [2, 7-9].

[0049] VGF is proteolytically processed with increasing differentiation of neuronal cells via neuron-specifically expressed endoproteases, which presumably recognize basic amino acids. As Trani et al. were able to show, C-terminal VGF peptides with masses of 20, 18 and 10 kDa are produced [10]. This VGF processing takes place in the postendoplasmic reticulum. These peptides accumulate in secretory vesicles, are released preferably by membrane depolymerization and might possibly play a role in neuronal communications [10]. Prohormone convertases such as, for example, PC1, PC2 or PC3 are known from the literature as examples of endoproteases which proteolytically cleave protein precursor molecules at dibasic sequence sites. The VGFARP peptides identified by us are, bowever, surprisingly fragments with a distinctly lower molecular weight than 10 to 20 kDa, and are therefore different from the VGF peptides described by Trani et al. In addition, the anti-VGF antibodies used by Trani et al. to detect these VGF peptides recognize VGFARP peptides which are different from the sequences of the VGFARP peptides. We have detected VGFARP peptides both in Alzheimer's disease patients and in the control group. The peptides identified by us represent novel VGF processing products which have not previously been described. The concentrations of the VGFARP neptides may be either uniformly raised or else uniformly lowered, in a manner which is specific for each peptide, in the patient group relative to the control group. Exclusively other VGF peptides of unknown sequence, derived from the C-terminal region of the VGF protein and having a distinctly higher molecular weight than the peptides newly identified and sequenced for the first time by us, were previously known [10]

PREFERED EMBODIMENTS OF THE INVENTION

[0950] The chronic dementia disease detected by the method of the invention is preferably Abbeimer's disease. It has been possible to date to detect the change in the concentration of the peptides and peptide fragments in the invention in Alzbeimer's disease patients. It can be concluded from this that the peptides of the invention can be used for the detection and for the therapy of Alzbeimer's disease and related purpological diseases.

[9051] The identification is prefurably concentrated on particular peptide fragments of the VGF probins baving the GeneBank accession No. NM_003378, or the DDBI accession No. Y12561 (Seq. IDS 43 and 44), i.e. on peptides which comprise partial sequences of these VGF proteins. These VGF peptides (VGF protein fragments) are referred to as VGF derived Abbeliumer related peptide (VGFARP) and they are represented by Seq. ID 1 to 42. The alignment of the VGF proteins and VGFARP peptides is depicted in Fig. 1. The sequences we found for the peptides are indicated in the sequence issing a

[9052] We have detected various VGF peptides derived from two VGF protice variants for the first time in biological samples. These peptides, which are referred to as VGFARP peptides, perspected defined fragments of VGF proteins. These fragments are produced in a natural way in nature and have not previously been described in the literature. These fragments are different from peptides generated in the literature often by in vitor proteolysis by addition of proteases such as, for example, trypsin). They therefore represent more, proviously unknown substances. These peptides were initially enriched and partified from biological samples by what systemetry from other accompanying peptides, so that it was subsequently possible to sequence these VGFARP periodics.

TABLE 1

single-letter amino acid code are as follows:							
	Sequenz sition	VGFARP	Seq.	Monoisotop theoret. mass (Da)	Sequence		
Y12661 23-59	NM_003378 23-59	1	1	3666.8278	APPGRPEAQPPPLSSEH KEPVAGDAVPGPKDGSA PEV		
23-62	23-62	2	2	3950.9875	appgrpeaqppplsseh kepvagdavpgpkdgsa pevrga		
23-58	23-58	18	15	3567.7594	appgrpeaqppplsseh Kepvagdavpgpkdgsa PE		
24–59	24-59	3	3	3595.7907	PPGRPEAQPPPLSSEHK EPVAGDAVPGPKDGSAP EV		

The sequences of the peptides in the

TABLE 1-continued

The sequences of the peptides in the single-letter amino acid code are as follows:

Monoisotop

VGF-S Posi	equenz ition	VGFARP No.	Seq.	theoret. mass (Da)	Sequence
24-62	24-52	4	4	3879.9504	PPGRPEAQPPPLSSEHK EPVAGDAVPGPKDGSAP EVRGA
26-59	26-59	5	5	3401.6852	GRPEAQPPPLSSEHKEP VAGDAVPGPKDGSAPEV
26-61	26-61	6	6	3614.8077	GRPEAQPPPLSSEHKEP VAGDAVPGPKDGSAPEV RG
26-62	26-62	7	7	3685.8448	GRPEAQPPPLS SEHKEP VAGDAVPGPKDGS APEV RGA
26-58	25-58	19	16	3302.6167	GRPEAQPPPLSSEHKEP VAGDAVPGPKDGSAPE
26-57	26-57	20	17	3173.5741	GRPEAQPPPLSSEHKEP VAGDAVPGPKDGSAP
26-64	25-64	21	18	3955.9889	GRPEAQPPPLSSEHKEP VAGDAVPGPKDGSAPEV RGARN
49-62	49-62	10	8	1336.6735	PGPKDGSAPEVRGA
90-114	90-114	22	19	2503.1827	LDRPASPPAPSGSQQGP EEEAAEAL
*50 _{r1} 57 _{+r2}	50 _{+r1} -57 _{+r3}	15	12	≧727.3501	r1-GPKDGSAP-r2
39-46	39-46	23	20	851.4137	r7-HKEPVAGD-r8
50-57	98-105	24	21	≧730.3246	r9-APSGSQQG-r10
-	121-156	25	22	3745.7343	SQTHSLPAPESPEPAAP PRPQTPENGPEASDPSE EL
164-174	164-174	26	23	1235.5782	QELRDFSPSSA
133 _{+r11} 140 _{+r12}	133 _{+r11} - 140 _{+r12}	27	24	≧833.4395	rll-EPAAPPRP-rl2
351-418	-	11	9	7518.2744	LQEAABERESAREEBA EQERROGEERVGEBDE AABAABABABABABARARQ NALLFABBBDGBAGABD
350-367	350-367	28	25	2031.8981	GLQEAAEERESAREEEE A
350-370	350-370	29	26	2418.0419	glqeaaeeresareeee aeqe
-	373-417	30	27	4806.0408	GGEERVGEEDEEAAEAE AEAEEAERARQNALLFA EEEDGEAGAED
-	373-404	31	28	3456.5513	GGEERVGEEDEEAAEAE AEAEEAERARQNALL
374-418	-	32	29	4806.0408	GEERVGBEDERAARAE AEADEAERARQNALLFA EEEDGEAGAED

TABLE 1-continued

The	sequences	of the	peptides	in t	he
single-l	etter amin	acid	code are	as fe	ollows:

_					
UCFc.	equenz	VGFARP	Sen	Monoisotop theoret.	
Posi	No.	ID	mass (Da)	Sequence	
421-456	420-455	33	30	4058.7043	SQEETPGHRRKEAEGTE EGGEEEDDEEMDPQTID SL
++ 421–472	420-471	12	10	5776.6294	SQEETPGHRRKEAEGTE EGGEEEDDEEMDPQTID SLIELSTKLHLPADDVV S
421-479	420-478	13	11	6618.0363	SQEETPGHRRKEAEGTE EGGEEEDDEEMDPQTID SLIELSTKUHLPADDVV SIIEEVEE
460-472	459-471	34	31	1380.7249	STKLHLPADDVVS
355 _{+r13} - 362 _{+r14}	355 _{+r13} - 362 _{+r14}	35	32	≥946.4468	r13-AEERESAR-r14
381 ₊₂₃ 389 ₊₂₄	381 ₊₂₃ 389 ₊₂₄	16	13	≧862.3192	r3-EDEEAAEA-r4
446 _{+e5} - 453 _{+x6}	445 ₊₂₅ - 452 ₊₂₅	17	14	≧961.4063	r5-EEMDPQTI-r6
-	485-522	36	33	3903.0180	nappepupppraadapt Hurspopppprapapard Elpd
-	485-521	37	34	3787.9911	NADPEPUPPPRAADAPT HVRSPQPPPPAPAPARD ELP
501 _{+r15} - 508 _{+r16}	500 _{+r15} - 507 _{+r16}	38	35	≧920,4828	r15-PTHVRSPQ-r16
26-31	26-31	39	36	656.3242	GRPEAQ
25-62	25-62	40	37	3782.8976	PGRPEAQPPPLSSEHKE PVAGDAVPGPKDGSAPE VRGA
177-193	177—193	41	38	1886.8970	QQETAAAETETRTHTLT
177-191	177-191	42	39	1672.7653	QQETAAAETETRTHTLT
180-187	180-187	43	40	≧792.3501	r17-taaaetet-r18
374-404	-	44	41	3343.4672	GEERVGEEDEEAAEAAE AEADEAERARQNAL
457-476	456-475	45	42	2220.1889	IELSTKLHLPADDVVSI IEE
¥126	il - Prote	sin	43	deduced	GP-protein sequence from Y12661 of the ta Bank of Japan
NM_003	378 - Pro	tein	44		/FG-protein sequence rom NM_003378 of the

TABLE 1-continued

The sequences of the peptides in the single-letter amino scid code are as follows:

VGF-Sequenz Position	VGFARP No.	Seq.	Monoisotop theoret. maaa (Da) Sequence
Y12661 - DNA		45	Complete VGF-DNA sequence from ENA Data Bank of Japan
NM_003378 - DY	IA	46	Complete VGF-DNA sequence from NCBI Bata Bank

^{*} rl represents a sequence which corresponds to the sequence or parts of the sequence of the VGF protein from smino scid 49-23, and r1 can be between 0 and 27 amino acids long, atarting from amino acid 50 of the VGF protein. Correspond ingly, r2 represents the VGF protein sequence from amino acid 58 to 64 or parts thereof, and r2 can be between 0 and 7 amino acida long, atarting from VGF amino acid 57. r3 represents the VGF protein sequence from amino acid 380 to 373 or parts therof, r4 represents the VGF protein acquence from amino acid 389 to 418 or parts thereof, r5 represents the WGF asquence from amino acid 445 to 421 or parts thereof, r6 representa the VGF protein sequence from amino acid 454 to 479 of parts thereof, r7 represents the VGF protein sequence from amino acid 38 to 23 of parts thereof, r8 represents the VGF protein sequence from amino acid 47 to 64 or parts thereof, r9 represents the VGF protein sequence from amino acid 97 to 90 or parts thereof, r10 represents the VGF protein acquence from amino acid 106 to 114 or parts thereof, r11 represents the VGF protein sequence from amino acid 132 to 121 or parta thereof, r12 represents the VGF protein sequence from amino acid 141 to 156 or parts thereof, r13 represents the VGF protein sequence from amino acid 354 to 350 or parts thereof r14 represents the VGF protein sequence from amino acid 363 to 370 or parts thereof, r15 represents the VGP protein sequence from amino acid 500 to 486 or parts thereof, r16 represents the VGF protein sequence from amino acid 509 to 523 or parts thereof, r17 represents the VGF protein sequence from amino acid 179 to 177 or parts thereof, r18 represents the VGF protein sequence from amino acid 192 to 193 or parts thereof. VGFARP-12 was identified as nonoxidized and as monooxidized

peptide (increase in the molecular weight by about 16 dal-

[0053] Suitable Peptides

[0054] The peptides can exist in post-translational or chemical modification forms, thus influencing inter this their masses and the identification by mass spectrometry and also the clustion behavior on chromatography such as, for example, on reverse phase chromatography. In particular, the peptides may be in glycosylsted, phosphorylated, sail-fated, amidated, oxidized etc. form in the sample to be investigated. The modified peptides are preferably in the form of peptides oxide such as, for example, the peptide VGFARP-12 which was identified both as unmodified peptide and as peptide oxide.

[0055] The peptides are also regarded as VGRARP peptides in particular when individual amino acids differ from the corresponding sequence of the VGR protein, in particular when a maximum of 2 amino acids differ from the VGP protein sequence. It is permissible in this connection for there to be point mutations, deletions, internal insertions of amino acids, and N- and C-terminal extensions, as long as the VGFARP peptide sequence comprises at least 8 amino acids which are conserved, i.e. unchanged, relative to the amino acid sequence of the relevant VGF protein. VGFARP 3P corresponds no acception, as in other vocations 6 amino acids. [905G] For a positive desection of the disease, it is furthermore provided in a further development of the investion for the concentration of the identified peptide(s) to be raise or lowend for each of these peptides in a specific manurelative to the concentration of the respective peptide in a control sample. The ratio of the concentrations of the respective peptides to the concentration of the control sample can be used to determine the severity of the disease.

[0057] The control sample may be a pooled sample from various controls. The sample to be investigated may also be a pooled sample, and where there is a positive result individual investigations are subsequently carried out.

[0058] Suitable Biological Samples

[9059] The biological sample may preferably be correspond that (CSF) or a sample such as serum, plasma, urice, stool, lear fluid, sprovial fluid, sputrum etc. This depends intert also not be sensitively of the chosen detection method (mass spectrometry, ELISA etc.). It is also possible where appropriate to use homogenized lissue samples, lissue sections and biopsy specimens. It is therefore provided in a further embodiment of this invention for tissue homogenates to be produced, for example from human tissue samples.

obtained in biopsies, for preparation of the sample to he investigated. These tissues can be commituted for expelle with manual homogenizers, with ultrasound homogenizers or with electrically operated homogenizers such as, for example, Ultramrax, and then be boiled in a manner known to the skilled worker in actific aqueous solutions with, for example, 0.1 to 0.2 M acetie acid for 10 minutes. The extracts are then subjected to the respective detection method, e.g. a mass spectrometric investigation. The samples can be prepared, for example where appropriate diluted or concentrated, and stored in the usual way.

[0060] Use of the VGFARP Peptides for Producing Diagnostic Agents

[0061] The invention further comprises the use of at least one VGRARP peptide of the invention or a 1 vGP protein for the diagnosis of neurological diseases, especially drongic dementia diseases, especially drongic dementia diseases, specially drongic dementia diseases, specially drongic dementia diseases, specially drongic antibodies or other agents which, because of their VGFARP peptides specific hindring properties, no saitable for developing diagnostic reagents for detecting these diseases. The invention also encompasses the use of VGFARP peptides for obtaining phage particles which hind these peptides specifically, or which conversely present VGFARP peptides on their surface and thus make it possible to identify hindring partners such as, for example, receptors of VGF proteins or VGFARP peptides.

[0062] Detection Methods for the VGFARP Peptides

[9063] Various methods can he used for detecting the VGPARP perigibles within the framework of the invention. Methods suitable are those which make it possible to detect VGPARP perigibles specifically in a patient's sample. Suitable methods are, inter alia, physical methods such as, for example, mass specimenty or liquid chomatography, molecular biology methods such as, for example, reverse transcriptuse polymerase chair nection (RT-PCR) or immunological detection techniques such as, for example, reuzyme linked immunosorbent assays (ELSA-A).

[0064] Physical Detection Methods

[9065] One embodiment of the invextion is the use of physical methods which are able to indicate the peptides of the invention qualitatively or quantitatively. These methods of the invention qualitatively or quantitatively. These methods include, inter tall, mass spectrometry, liquid chromatography, thin-layer chromatography, NMR (unclear magnetic play, thin-layer chromatography, NMR (unclear magnetic of quantitative measured results from a sample to he investigated with the measurements of thinted in a group of presidents suffering from neurological diseases, in particular knowled demental diseases, preferably Alzbeimer's disease, and a control group. It is possible to infor the presence of a neurological diseases, in particular a chronic demental disease, in particular Alzbeimer's disease, and of the severity of this disease from these results.

[9066] In a preferred embodiment of this invention, the peptides in the sample are separated by chromotography before the identification, in particular preferably by reverse phase chromotography, with particular preferably by reverse phase chromotography, with particular preferable for separation of the peptides in the sample by high-resolution reverse phase high performance chromotography (RP-HPLC). A further embodiment of this invention is the carrying out of precipitation reactions to fractionate the sample using precipitants such as, for example, aumonium sulfate, polyedhyfone dyvod, trichbrocaetic acid, acotone, chanol etc. The fractions oltatined in this way are subjected singly to the respective discretion method, e.g. the investigation using mass spectrometry. A further embodiment of the invention is the use of liquid phase extraction. For this purpose, the sample is mixed with a mixture of an organic solvent such as, for example, polyethylene glycol (PEG) and an aqueous salt solution. Owing to their physical properties, particular constituents of the sample then accumulate in the organic phase, and others in the aqueous phase, and can thus be separated from one another and subsequently analyzed further.

[0067] Reverse Phase Chromatography

[0068] A particularly preferred embodiment of this invention encompasses the use of reverse phase chromatography, in particular a C18 reverse phase chromatography column using mobile phases consisting of trifluoroacetic acid and acetonitrile, for separation of peptides in human cerebrospinal fluid. For example the fractions collected in each case each comprise 1/100 of the mobile phase volume used. The fractions obtained in this way are analyzed with the aid of a MALDI mass spectrometer (matrix-assisted laser desorption ionization) using a matix solution consisting of, for example, of L(-) fucose and alpha-cyano-4-hydroxycinnamic acid dissolved in a mixture of acetonitrile, water, trifluoroacetic acid and acetone, and thus the presence of particular masses is established and the signal intensity quantified. These masses correspond to the masses of the VGFARP peptides of the invention.

[0069] Mass Spectrometry

[0070] In a preferred emhodiment of the invention, VGFARP peptides can be identified with the aid of mass spectrometric determination, preferably a MALDI (matrixassisted laser desorption and ionization) mass spectrometry. In this case, the mass spectrometric determination further preferably includes at least one of the following mass signals, in each case calculated on the hasis of the theoretical monoisotopic mass of the corresponding peptide. It is possible for slight differences from the theoretical monoisotopic mass to show owing to the experimental error and the natural isotope distribution. In addition, in MALDI mass determinations a proton is added to the peptides owing to the method of measurement, whereby the mass increases by 1 dalton. The following masses correspond to the theoretical monoisotopic masses of the peptides identified by us; calculated with suitable software, in this case GPMAW 4.02. These theoretical monoisotopic masses may occur singly or in combination in a sample: VGFARP-1 (SEQ ID NO:1) =3666.8278/VGFARP-2 (SEQ ID NO:2)=3950.9875/VG-FARP-18 (SEQ ID NO:15)=3567.7594/VGFARP-3 (SEQ ID NO:3)=3595,7907/VGFARP-4 (SEO ID NO:4)= 3879.9504/VGFARP-5 (SEQ ID NO:5)=3401.6852/VG-FARP-6 (SEQ ID NO:6)=3614.8077/VGFARP-7 (SEQ ID NO:7)=3685.8448/VGFARP-19 (SEQ ID NO:16)= 3302.6167/VGFARP-20 (SEQ ID NO:17)=3173.5741/VG-FARP-21 (SEO ID NO:18)=3955.9889/VGFARP-10 (SEO ID NO:8)=1336.6735/VGFARP-22 (SEQ ID NO:19)= 2503.1827/VGEARP-15 (SEO ID NO:12)=≥727.3501/VG-FARP-23 (SEQ ID NO:20)-≥851.4137/VGFARP-24 (SEQ ID NO:21)=≥730.3246/VGFARP-25 (SEO ID NO:22)= 3745.7343/VGFARP-26 (SEQ ID NO:23)=1235.5782/VG- FARP-27 (SEQ ID NO:24)=≥833.4395/VGFARP-11 (SEQ ID NO:9)=7518.2744/VGFARP-28 (SEQ ID NO:25)= 2031.8981/VGFARP-29 (SEO ID NO:26)=2418.0419/VG-FARP-30 (SEQ ID NO:27) 4806.0408/VGFARP-31 (SEQ ID NO:28)=3456.5513/VGFARP-32 (SEQ ID NO:29)= 4806.0408/VGFARP-33 (SEQ ID NO:30)=4058.7043/VG-FARP-12 (SEO 1D NO:10)=5776.6294/VGFARP-13 (SEO ID NO:11)=6618.0363/VGFARP-34 (SEO ID NO:31)= 1380.7249/VGFARP-35 (SEQ ID NO:32)=≥946.4468/VG-FARP-16 (SEO ID NO:13)=≥862.3192/VGFARP-17 (SEO ID NO:14)=≥961.4063/VGFARP-36 (SEQ ID NO:33)= 3903.0180/VGFARP-37 (SEO ID NO:34)=3787,9911/VG-FARP-38 (SEQ ID NO:35)=≥920.4828/VGFARP-39 (SEQ ID NO:36)=656.3242/VGFARP-40 (SEQ ID NO:37)= 3782.8976/VGFARP-41 (SEO ID NO:38)=1886.8970/VG-FARP-42 (SEQ ID NO:39)=1672.7653/VGFARP-43 (SEQ ID NO:40)=≥792.3501/VGFARP-44 (SEO ID NO:41)= 3343.4672 and VGFARP-45 (SEO ID NO:42)=2220.1889.

[9071] The symbol—E (is greater than or equal to) is to be understood to mean that the relevant VGFARP peptides cannot have any larger masses but can have only the masses possible owing to the amino acids which are possibly additionally present at the eads of these peptides. Amino acids which may be additionally present at the eads of these peptides are not just any once but only those which may be prises at a this sequence position owing to the sequence of the VGF protein.

[0072] Mass Spectrometric Determination of the Sequence of the VGFARP Peptides

[6073] For the further practical application of this embodiment, further confirmation of the result of election is advisable and possible by establishing the identity of the peptides corresponding to the masses, taking account exsistely of peptide signals which may be derived from a VGF proton. This confirmation takes place by identifying the peptide signals preferably using methods of mass spectromerty, e.g. MS/MS analysis [11].

[0074] Novel, specific peptides of VGF proteins (VGFARP peptides) were identified, and their significance was revealed by the method of the invention. These peptides and their derivatives are referred to herein as VGFARP peptides. Their sequences are indicated in the sequence listing. The VGFARP peptides VGFARP-15 (SEQ ID NO:12), 16 (SEQ ID NO:13), -17 (SEQ ID NO:14), -27 (SEQ ID NO:24), -35 (SEQ ID NO:32), 38 (SEQ ID NO:35) and VGFARP-43 (SEO ID NO:40) may comprise on the Nand/or C-terminus additional amino acids corresponding to the corresponding sequence of the relevant VGF protein. The invention also encompasses the VGFARP peptides prepared recombinantly or synthetically, and isolated from biological samples, in unmodified, chemically modified or post-translationally modified form. In this connection, two point mutations and other differences are possible as long as the VGFARP peptide has at least 8 amino acids which agree in their identity and their position within the peptide sequence with a VGF protein.

[0075] Molecular Biology Detection Techniques

[0076] Finally, the invention also encompasses nucleic acids which correspond to VGFARP peptides, and especially those which correspond to the VGFARP peptides of the invention, the use thereof for the indirect determination and

quantification of the relevant VGF proteins and peptides. This also includes uncleic acids which represent for example, noncoding sequences such as, for example, S- or 3-untranslated regions of the mRNA, or nucleic acids which show a sequence agreement with the VGF nucleic acid sequence which is sufficient for specific hybridization experiments and which are therefore suitable for the indirect detection of relevant proteins, especially the VGFARP peptides.

[0077] One exemplary embodiment thereof encompasses the obtaining of tissue samples, e.g. of biopsy specimens, from patients and the subsequent determination of the concentration of an RNA transcript corresponding to the gene having the GeneBank accession No. NM_03378 or the accession No. Y12661 of the DNA Data Bank of Japan. DDBJ or corresponding to homologous VGF variants. This entails comparison of quantitative measured results (intensities) from a sample to be investigated with the measurements obtained in a group of patients suffering from Alzheimer's disease and a control group. Methods which can be used for the quantification are, for example, reverse transcriptase polymerase chain reaction (RT-PCR), quantitative real-time PCR (ABI PRISM® 7700 Sequence Detection System, Applied Biosystems, Foster City, Calif., USA), in situ hybridization or Northern blots in a manner known to the skilled worker. The presence of a chronic dementia disease, preferably Alzheimer's disease and/or the severity thereof can be inferred from the results.

[0078] Immunological Detection Methods

[0079] In a further preferred embodiment of the invention, the VGFARP peptides or the VGF proteins can be identified using an immunological detection system, preferably an ELISA (enzyme linked immuno sorbent assay). This immunological detection picks up at least one VGFARP peptide or VGF protein. To increase the specificity, it is also possible and preferred to use the so-called sandwich ELISA in which the detection of the VGFARP peptides depends on the specificity of two antibodies which recognize different epitopes within the same molecule. However, it is also possible to use other ELISA systems, e.g. direct or competitive ELISA, to detect VGFARP peptides or VGF proteins. Other ELISA-like detection techniques such as, for example, RIA (radio immuno assay), EIA (enzyme immuno assay), ELI-Spot etc. are also suitable as immunological detection systems. VGFARP peptides or VGF proteins isolated from biological samples, recombinantly prepared or chemically synthesized can be used as standard for the quantification. Identification of the VGFARP peptide(s) is generally possible for example with the aid of an antibody directed to the VGFARP peptide or VGF protein. Further methods suitable for such detections are, inter alia, Western blotting, immunoprecipitation, Dot-Blots, plasmon resonance spectrometry (BIACORE®-Technologie, Biacore International AB, Uppsala, Sweden), phage particles, PNAs (peptide nucleic acids), affinity matrices (e.g. ABICAP-Technologie, ABION Gesellschaft für Biowissenschaften und Technik mbH, Jülich, Germany) etc. Substances/molecules suitable as detection agents are generally all those permitting the construction of a specific detection system because they specifically bind a VGFARP peptide of VGF [0080] Obtaining of VGFARP Peptides and Anti-VG-FARP Peptide Antibodies

[0081] A further embodiment of the invention is the obtaining of VGFARP peptides using recombinant expression systems, chromatographic methods and chemical synthesis protocols which are known to the skilled worker. The VGFARP peptides obtained in this way can be used inter alia as standards for quantifying the respective VGFARP peptides or as antigen for producing VGFARP peptide antibodies. Methods known to the skilled worker and suitable for isolating and obtaining VGFARP peptides include the recombinant expression of peptides. It is possible to use for the expression of the VGFARP peptides inter alia cell systems such as, for example, bacteria such as Escherichia coli, yeast cells such as Saccharomyces cerevisiae, insect cells such as, for example, Spodoptera frugiperda (Sf-9) cells, or mammalian cells such as Chinese Hamster Ovary (CHO) cells. These cells are obtainable from the American Tissue Culture Collection (ATCC). For recombinant expression of VGFARP peptides, for example nucleic acid sequences which code for VGFARP peptides are inserted in combination with suitable regulatory nucleic acid sequences such as, for example, promoters, antihiotic selection markers etc. into an expression vector by molecular hiology methods. A vector suitable for this purpose is, for example, the vector pcDNA3.1 from Invitrogen. The VGFARP peptide expression vectors obtained in this way can then be inserted into suitable cells, e.g. by electroporation. The VGFARP pentides produced in this way may be C- or N-terminally fused to heterologous sequences of peptides such as polyhistidine sequences, hemagglutinin epitopes (HAtag), or proteins such as, for example, maltose-hinding proteins, glutathione S-transferase (GST), or protein domains such as the GAL-4 DNA binding domain or the GAL4 activation domain. The VGFARP peptides can he prepared by chemical synthesis for example in accordance with the Merrifield solid-phase synthesis protocol using automatic synthesizers which are obtainable from various manufacturers.

[0082] A further embodiment of this invention is the isolation of VGFARP peptides from hiological samples or cell culture media or cell lysates from recombinant expression systems, e.g. using reverse phase chromatography, affinity chromatography, ion exchange chromatography, gel filtration, isoelectric focusing, or using other methods such as preparative immunoprecipitation, ammonium sulfate precipitation, extraction with organic solvents etc. A further embodiment of the invention is the obtaining of monoclonal or polyclonal antibodies using VGFARP peptides. The obtaining of antibodies takes place in the conventional way familiar to the skilled worker. A preferred emhodiment of the production and ohtaining of VGFARP peptide-specific antibodies, and a particularly preferred embodiment is the production of VGFARP peptide-specific antibodies which recognize necepitopes, i.e. epitopes which are present only on VGFARP peptides but not in a VGF protein. Such anti-VGFARP peptide antibodies make the specific immunological detection of VGFARP peptides possible in the presence of VGF protein. Polyclonal antihodies can be produced by immunizations or experimental animals such as, for example, mice, rats, rabbits or goats. Monoclonal antibodies can be obtained for example by immunizations of experimental animals and subsequent application of hybridoma techniques or else via recombinant experimental approaches such as, for example, via antihody libraries such as the HuCAL® autibody library of MonphoSys, Martinsried, Germany, or other recombinant production methods known to the skilled worker. Autibodies can also be used in the form of autibody fragments such as, for example, Fah fragments or Fab2 fragments etc.

[0083] Therapy Development and Monitoring Through VGFARP Peptide Determinations

[0084] A further exemplary use is the quantitative or qualitative determination of the abovementioned VGFARP peptides or VGF proteins for estimating the efficacy of a therapy under development for neurological diseases, in particular chronic dementia diseases, in particular Alzheimer's disease. The invention can also he used to identify suitable patients for clinical studies for developing therapies for these diseases, in particular Alzheimer's disease. This entails comparison of quantitative measured results from a sample to he investigated with the measurements obtained in a control group and a group of patients. The efficacy of a therapeutic agent, or the suitability of the patient for a clinical study, can be inferred from these results. The testing of efficacy and the selection of the correct patients for therapies and for clinical studies is of outstanding importance for successful application and development of a therapeutic agent, and no clinically measurable parameter making this reliably possible is yet available for Alzheimer's disease

[9085] Examination of the Therapeutic Efficacy of VGF Proteins, VGFARP Peptides and of Agents which Modulate the Expression and the Bioavailability of these Substances

[0086] One exemplary embodiment thereof encompasses the cultivation of cell lines and their treatment with VGF proteins, VGFARP peptides or with substances which promote the expression of VGF protein, such as, for example, NGF, BNDF or NT-3, or promote the processing of VGF protein to VGFARP pentides, such as, for example, prohormone convertases. It is possible thereby to establish the biological properties of VGF protein and VGFARP peptides in connection with neurological diseases, in particular Alzheimer's disease. Fusion proteins and fusion peptides can also be used for the treatment of the cell lines, e.g. fusion proteins consisting of prohormone convertases fused to peptide sequences which promote transport of the fusion protein into the interior of the cell. Examples of possible fusion partners of, for example, prohormone convertases are HIV TAT sequences or antennapedia sequences etc. It is likewise possible to transfect cell lines with expression vectors which bring about, directly or indirectly, expression of VGF protein or VGFARP peptides by the transfected cells. These expression vectors may code inter alia for VGFARP peptides, VGF proteins, NGF, BNDF, NT-3 or for prohormone convertases. Transfection of combinations of the said proteins can also he carried out. Alternatively. suitable cell lines can be treated with anti-VGF protein or anti-VGFARP peptide antibodies or with nucleic acids which suppress the expression of VGF, such as, for example, VGF antisense nucleic acids, VGF triplex nucleic acids or ribozymes directed against VGF mRNA. Treatment with anti-NGF, anti-BNDF or anti-NT-3 antihodies might also be carried out to suppress VGF protein expression. Cell lines which appear suitable as neurological model systems in connection with VGF in particular can be used for such investigations. Read-out systems which can be used for

these investigations are inter alia tests which measure the rate of proliferation of the treated cells, their metabolic activity, the rate of apoptosis of the cells, changes in cell morphology, in the expression of cell-intrinsic proteins or reporter genes or which measure the release of cytosolic cell constituents as markers for cell death. Further test systems which can be used are suitable strains of experimental animals, e.g. of mice or rats, which are considered as model of neurological diseases, in particular as model of Alzheimer's disease. These experimental animals can be used to investigate the efficacy of therapentic strategies which aim to modulate the concentration of VGFARP peptides or of VGF proteins. It is additionally possible to investigate proteins and peptides such as, for example, VGF proteins, VGFARP peptides, NGF, BNDF, NT-3, prohormone convertases etc. in experimental animals, it being possible for these peptides and proteins in some circumstances to be pharmaceutically processed so that they are better able to cross the blood-brain barrier and/or the blood-CSF barrier. It is possible to use as pharmaceutical processing method inter alia liposome-packaged proteins and peptides, proteins and peptides fused to transport sequences such as, for example, an HIV TAT sequence etc. In addition, peptides and proteins can be chemically modified in such a way that they acquire more lipophilic properties and are therefore able to penetrate more easily into cells. Peptides which are only slightly soluble in aqueous solutions can conversely be chemically modified so that they become more hydrophilic and then can be used for example as intravenously injectable therapeutic agent. Acid-resistant capsules can be used to protect sensitive substances, intended for oral administration, in the stomach.

[0087] Read-out parameters in experiments with animal models may be the survival time of the animals, their behavior and their short-term memory. One example of a memory test which is suitable for experimental animals is be used are the determination of body function such as, for example, blood tests, measurement of brain currents, metabolism test, the rate of expression of Voff protein and VOFARP peptides and other preteins associated with the disease, and morphological and histological investigations on itssues such as, for example, the brain.

[0088] Methods of Treatment

[0089] Another embodiment of the invention comprises methods of treatment of neutrological diseases, in particular of chronic dementia diseases, like Alheimer disease, etc. At least noe of the peptides, nucleic acids, autitodies, agonists or antagonists as defined herein may be used therein. The method may result in a reduction or increase, respectively, in the concentration of the altered VGFARP peptides or VGF proteins.

[0900] In particular, the method comprises administering abanthodies directed against Voff proteins, VOFfARP peptides, NGR, BNDF or NT-3 are administered, and/or by autiense medice acids, tripbæ medice acids or ribozymes are administered, in order to reduce the expression of Voff proteins, VOFFARP peptides, NGFARP peptides or VGFARP peptides.

peptides. Alternatively, the method comprises administering to a patient suffering from a neurological disease for an increase of the concentration of VGFARP peptides a) VGF proteins, VGFARP peptides, NGF, BNDF or NF-3, and/or b) mucleic acids which code for VGF proteins, VGFARP peptides, NGF, BNDF or NF-3, and/or c) substances which protein the proteins, and/or d) agonitis of the VGFARP peptides or of VGF proteins and/or d) agonitis of the VGFARP peptides or of VGF proteins are administered to a patient.

[0091] The invention is illustrated in detail below by means of examples. Reference is also made to the figures in this connection.

[9092] FIG. 1 shows an alignment of the peptides of the invention with two known variants of the VGP point which are identified in the figure by their database accession No. NM. 003376 (SBQ II) NO.44) and Y1264 (SBQ II) NO.44). Sequence positions which are identical in both variants of the VGP proteins are represented by an asterisk in the sequence of NM. 003378 (SEQ II) NO.44). Different sequences are represented by the amino acid code in white letters on black background. The arrow at the end or at the start of partial sequences of VGFARP-12 (SEQ II) NO.13), 45 (SEQ II) NO.42) and 34 (SEQ II) NO.31) indicates that the respective sequence extends over two lines in the alignment.

[0093] FIG. 2 shows a chromatogram recorded using reverse phase chromatography as in Example 2 for the separation and enrichment of the VGF peptides from cerebrospinal fluid.

[0094] FIG. 3 shows a spectrum resulting from MALD in mass spectrometric measurement as in Example 3 of VGRARP.7 (SEO ID NO.7), with a theoretical monistopic mass of 3605 dallon, after reverse phase chromatography of human cerebrospinal fluid as in Example 2. VGRARP.7 (SEQ ID NO.7) corresponds to the VGRARP.4 (SEQ ID NO.7) corresponds to the VGRARP.4 (Sequence of Seq. ID 43 (accession No. Y1266)) of amino acid 26-62.

[9095] FIG. 4 shows data generated by MALDI as relatively quantifying MS method. A sample was mixed involved quantifying MS method. A sample was mixed intensity both of these standard spenials, and the intensity both of these standards signals and of representative sample signals was measured. All signal intensities of the standards were standardized to their signal intensit of the at concentration of 0.54 M/ (-1). Each peptide shows an individual typical ratio of signal strength to concentration, which can be read off in this diagram from the gradient of the plot.

[0096] FIG. 5 shows an MS/MS fragment spectrum as in Example 4 of the peptide VGFARP-13 (SEQ ID NO:11) of the invention

[0097] Upper trace: raw data of the measurement.

[0098] Lower trace: converted, deconvoluted mass spectrum of VGFARP-13.

[0099] The peak pattern is characteristic of VGFARP-13 (SEQ ID NO:11). VGFARP-13 (SEQ ID NO:11) corresponds to the VGF sequence of Seq. ID 43 (accession No. Y12661) of amino acid 421-479.

[0100] FIGS. 6A to 6C show in the form of box-whisker plots a comparison of the integrated MALDI mass spectrometric signal intensities of various VGFARP peptides in controls, compared with the signal intensities in samples from Alzheimer's disease patients.

EXAMPLE 1

Obtaining Cerebrospinal Fluid for Determining VGFARP Peptides

[0101] CSF or cerebrospinal fluid (fluid of the brain and spinal cord) is the fluid which is present in the four ventricles of the brain and in the substractmoid space and which is produced in particular in the choosid plexas of the lateral ventricle. Cerebrospinal fluid is usually taken by lembar purcture and less often by subsociptial puncture or ventricular puncture. In lumbar puncture (spinal paneture), to the eccetorogain thin, they neutron involves penetration of the spinal substractmoid space between the 3rd and 4th or the metalle, and thus CSF being observed. This long bollow metalle, and thus CSF being observed. The simple is centrifuged at 2000cg for 10 minutes, and the supernitatal is stored at 4-60° C.

EXAMPLE 2

Separation of Peptides in Cerebrospinal Fluid (CSF) for Mass Spectrometric Measurement of VGFARP Peptides

[0102] For the detection of VGF peptides in CSF by mass spectrometry, it is necessary in this example to separate the peptide constituents. This sample pretreatment serves to concentrate the peptides of the invention and to remove components which may interfere with the measurement. The separation method carried out is a reverse phase chromatography. Various RP cbromatography resins and eluants are equally suitable for this. The separation of VGF peptides using a C18 reverse phase chromatography column with the size of 4 mm×250 mm supplied by Vydac is [lacuna] by way of example below. Mobile phases of the following composition were used: mobile phase A: 0.06% (v/v) trifluoroacetic acid, mobile phase B: 0.05% (v/v) trifluoroacetic acid, 80% (v/v) acetonitrile. Chromatography took place at 33° C. using an HP ChemStation 1100 supplied by Agilent Tecbnologies with a micro flow cell supplied by Agilent Technologies. Human cerebrospinal fluid was used as sample. 440 µl of CSF were diluted with water to 1650 µl, the pH was adjusted to 2-3, the sample was centrifuged at 18 000x for 10 minutes and finally 1500 μ l of the sample prepared in this way were loaded onto the chromatography column. The chromatography conditions were as follows: 5% mobile phase B at time 0 min, from time 1 to 45 min continuous increase in the mobile phase B concentration to 50%, from time 45 to 49 min continuous increase in the mobile phase B concentration to 100% and subsequently up to time 53 min constant 100% buffer B. Collection of 96 fractions each of 0.5 ml starts 10 minutes after the start of the chromatography. The chromatogram of a cerebrospinal fluid sample prepared under the experimental conditions described berein is depicted in FIG. 2.

EXAMPLE 3

Measurement of Masses of Peptides by Means of MALDI Mass Spectrometry

[0103] For mass analysis, typical positive ion spectra of peptides were produced in a MALDI-TOF mass spectrometer (matrix-assisted laser desorption ionization). Suitable MALDI-TOF mass spectrometers are manufactured by Per-Septive Biosystems Framingham (Voyager-DE, Voyager-DE PRO or Voyager-DE STR) or by Bruker Daltonik Bremen (BIFLEX). The samples are prepared by mixing them with a matrix substance which typically consists of an organic acid. Typical matrix substances suitable for peptides are 3,5-dimethoxy-4-hydroxycinnamic acid, \alpha-cyano-4-hydroxycinnamic acid and 2,5-dihydroxybenzoic acid. A lyophilized equivalent obtained by reverse phase chromatograpby and corresponding to 500 µl of buman cerebrospinal fluid is used to measure the VGFARP peptides of the invention. The chromatographed sample is dissolved in 15 ul of a matrix solution. This matrix solution contains, for example, 10 g/l \(\alpha\)-cyano-4-hydroxycinnamic acid and 10 g/l LC-)fucose dissolved in a solvent mixture consisting of acetonitrile, water, trifluoroacetic acid and acetone in the ratio 49:49:1:1 by volume. 0.3 µl of this solution is transferred to a MALDI carrier plate, and the dried sample is analyzed in a Voyager-DE STR MALDI mass spectrometer from PerSeptive Biosystems. The measurement takes place in linear mode with delayed extraction TM. An example of a measurement of one of the VGFARP peptides of the invention is shown in FIG. 3.

[0104] The MALDI-TOF mass spectrometer can be employed to quantify peptides such as, for example, the VGFARP peptides of the invention if these peptides are present in a concentration which is within the dynamic measurement range of the mass spectrometer, thus avoiding detector saturation. This is the case for the measurement of the VGFARP peptides of the invention in cerebrospinal fluid at a CSF equivalent concentration of 33.3 µl per µl of matrix solution. There is a specific ratio between measured signal and concentration for each peptide, which means that the MALDI mass spectrometry can preferably be used for the relative quantification of peptides. This situation is depicted in FIG. 4. If various amounts of different standard pentides are added to a sample, it is possible to measure the intensity both of these standard signals and of the sample signals. FIG. 4 shows by way of example a MALDI measurement as relatively quantifying MS method. All signal intensities of the standards were standardized to their signal intensity at a concentration of 0.64 µM (=1). Each peptide shows an individual, typical ratio of signal strength to concentration, which can be read off from the gradient of the plot.

EXAMPLE 4

Mass Spectrometric Identification of the VGFARP Peptides

[0105] For quantification of the VGFARP peptides of the invention it is necessary to ensure that the mass signals to be analyzed of peptides in the fractions obtained by reverse phase chromatography of cerebrospinal fluid, as in Example 2, in fact relate to the VGFARP pentides of the invention.

[0106] The peptides of the invention are employed in these fractions for example using nanoSpryn-MS/MS [11]. This entails a VGFARP peptide ion in the mass spectrometer being selected in the mass spectrometer on the basis of its specific mrk/(mss/charge) value in a manner known to the skilled worker. This selected ion is then fragmented by supplying collisional energy with an impinging gas, e.g., belium or nitrogen, and the resulting VGFARP peptide.

fragments are detected in the mass spectrometer in an integrated analysis unit, and corresponding me' values are determined (principle of tandem mass spectrometry) [13]. The fragmentation behavior of peptides makes unambiguous discutification of the VGBARP peptides of the invention possible when the accuracy of mass is, for example, 50 pem by the use of computer-assisted search methods [14] in sequence databases into which the sequence of a VGF protein has been entered. In this specific case, the mass spectrometric analysis took place with a Quadrupol-ToO Instrument, QSBar-Delasr model from Applied Biosystems-Sciez, USA. Examples of MS/MS fragment spectra are shown in FIG. 5.

EXAMPLE 5

Mass Spectrometric Quantification of the VGFARP Peptides to Compare Their Relative Concentration in Control Samples Compared with Patients' Samples

[0107] A sample preparation as in Example 1 and 2 followed by a MALDI measurement of the VGFARP peptides of the invention as in Example 3 were carried out on 222 clinical samples, i.e. 82 control samples and 130 samples from patients suffering from Alzheimer's disease. Examples of MALDI signal intensities are depicted in the form of hox-whisker plots in FIGS. 6A to 6C. The boxwhisker plots depicted in FIG. 6 are based on measurements carried out in each case on 29 to 45 samples from Alzheimer's disease patients, and 13 to 44 control samples per experiment. A total of 4 experiments was carried out. The hox-whisker plots depicted make it possible to compare the integrated MALDI mass spectrometric signal intensities of various VGFARP peptides in controls with the MALDI signal intensities in samples from Alzheimer's disease patients. In these, the box, i.e. the columns in the diagrams in FIGS. 6A to 6C, in each case includes the range of MALDI signal intensities in which 50% of the respective MALDI signal intensities are to be found, and the lines starting from the hox and pointing upward and downward (whiskers) indicate the range in which in each case the 25% of measurements which show the highest signal intensities (upper quarter) are to be found, and in which the 25% of measurements which show the lowest signal intensities (lower quarter) are to be found. The full line in the columns indicates the median and the broken line in the columns indicates the mean.

[0108] The headings in this document are intended merely to provide structure to the text. They are not intended to limit or restrict the matters described. All the examples are intended to characterize the concept of the invention in more detail but are not intended to restrict the equivalence range of the invention.

We claim:

 A method for detecting a chronic dementia disease or a predisposition to a chronic dementia disease in a patient in need thereof, comprising the steps of

obtaining a biological sample from said patient,

determining a concentration of at least one VGF protein or VGFARP peptide in the biological sample, and comparing the concentration of the at least one VGF protein or VGFAP peptide in the holoigeal sample to the concentration of the same protein or peptide in a concentration of the same protein or peptide in a concentration of the VGF protein or VGFARP peptide in the biological sample compared to the concentration of the VGF protein or VGFARP peptide in the control sample is indicative of chronic dementia diseases or a predisposition to a chronic demential disease.

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2. The method of claim I, wherein the at least one VGF protein or VGFARP peptide is selected from the group consisting of SEQ ID NO-45; a business of SEQ ID NO-45; and which the amino acid sequence of the mutant differs by a maximum of 20% from the amino acid sequence of SEQ ID NO-45; a mutant of SEQ ID NO-45 in a maximum of 20% from the amino acid sequence of SEQ ID NO-45; a mutant of SEQ ID NO-46 in maximum of 20% from the amino set dis expenses of SEQ ID NO-45; a protein that represents a naturally occurring allele of a VGF protein; a peptide derivate derivate from a VGFARP peptide a vulcate derivate from a VGFARP peptide, a VGFARP peptide available that differs by a maximum of 20 amino acids from the corresponding unmutanted VGFARP peptide.

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3. The method of claim 2, wherein said VGFARP peptide is selected from the group consisting of SEQ ID NO.1, SEQ ID NO.2, SEQ ID NO.3, SEQ ID NO.4, SEQ ID NO.5, SEQ ID NO.10, SEQ

- The method of claim 2 wherein the at least one VGF protein or VGFARP peptide is chemically modified.
- The method of claim 2 wherein the at least one VGF protein or VGFARP peptide is post-translationally modified.
- 6. The method of claim 1 whorein said method is carried out in combination with other diagnostic methods for chronic dementia diseases to increase the sensitivity and/or specificity thereof.
- 7. The method of claim 1 wherein the dementia disease is selected from the group consisting of Alzheimer's disease or a related neurological disease; Lewy body dementia; and vascular dementia.
- 8. The method of claim 1 wherein for a positive detection of the disease the concentration of the at least one VGF protein or VGFARP peptide is raised or lowered relative to the concentration of the VGF protein or VGFARP peptide in a control sample.
- 9. The method of claim 1 wherein the method is used to determine a parameter selected from the group consisting of: the severity of the disease, prognosis of the course of the disease, diagnosis of preliminary stages of neurological diseases, and mild cognitive impairment (MCI).
- 10. The method of claim 1 wherein the hiological sample is selected from the group consisting of cerebrospinal fluid, serum, plasma, urine, synovial fluid, stool, tear fluid, sputem and a tissue homogenate.

- 11. The method of claim 1 wherein the at least one VGF protein or VGFARP peptide is identified by mass spectrom-
- 12. The method of claim 11, wherein identification of the at least one VOE protein or VOEA-RP peptide by mass spectroscopy includes the determination of at least one of the theoretical monoisolopic mass peaks selected from the group consisting of 3666-8278/95/0-9875/55/75/94/3595.7991/3595.7991/3595.7991/3595.9991/3567.7994/3595.9991/3567.3596.3692/91/3595.9991/3567.3592/91/3567.3592/91/3567.3592/91/3595.9991/3567.3592/91/3567.3592/91/3595.9991/3567.3592/91/3567.3592/91/3595.9991/3567.3592/91/3692.99
- 3787 9911/≥920.4828/656.3242/3782.8976/1886.8970/ 1672.7655/≥792.3501/3343.4672 and 2220.1889 dalton. 13. The method of claim 1, wherein the at least one VGF protein or VGFARP peptide is identified with an immunological test.
- 14. The method of claim 13, wherein said immunological test is selected from the group consisting of enzyme linked immuno sorbent assay (ELISA), a radioimmunoassay, and a Western blot.
- 15. The method of claim 13, wherein the at least one VGF protein or VGFARP peptide is identified using a substance that binds to the protein or peptide.
- 16. The method of claim 15, wherein the substance that binds to the at least one VGF protein or VGFARP peptide selected from the group consisting of an antibody, an antibody fragment, a phage particle, and an affinity matrix.
- 17. The method of claim 1 further comprising the step of chromatographically fractionating said biological sample prior to said determining step.
- 18. The method of claim 17 wherein said step of chromatographically fractionating said hiological sample is carried out using reverse phase chromatography or high resolution reverse phase chromatography.
- 19. The method of claim 1 further comprising the step of fractionating the biological sample by precipitation reactions or liquid phase separations prior to said determining step.
 20. The method of claim 1 wherein said step of determining the comprehensive of the comprehe
- mining is carried out using antibodies against at least one VGF protein or VGFARP peptide.

 21. The method of claim 1 wherein said step of deter-
- 21. The method of claim 1 wherein said step of determining is carried out by detection of nucleic acids encoding at least one VGF protein or VGFARP peptide.
- 22. The method of claim 21 wherein the detection of nucleic acids is carried out using Northern blots, reverse transcriptase PCR or quantitative PCR.
- 23. A method for diagnosing a neurological disease in a patient, comprising the step of
 - obtaining a biological sample from said patient,
 - determining a concentration of at least one VGF protein or VGFARP peptide in the hiological sample, and
 - comparing the concentration of the at least one VGF protein or VGFARP peptide in the hiological sample to the concentration of the same protein or peptide in a control sample, wherein a difference between the concentration of the VGF protein or VGFARP peptide in the biological sample compared to the concentration of the VGF protein or VGFARP peptide in the control sample is middled to the concentration of the VGF protein or VGFARP peptide in the control sample is indicative of a neurological disease.

- 24. The method of claim 23 wherein said step of determining is carried out using antihodies against at least one VGF protein or VGFARP pentide.
- 25. The method of claim 23 wherein said step of determining is carried out by detection of nucleic acids encoding at least one VGF protein or VGFARP peptide.
- 26. The method of claim 25 wherein the detection of nucleic acids is carried out by using Northern blots, reverse transcriptase PCR or quantitative PCR.
- 27. The method of claim 23, wherein the method is used to monitor the efficacy of a therapy for a neurological disease.
- 28. The method of claim 23, wherein the method is used for stratifying patients who are suitable for therapies or clinical studies of neurological diseases.
- 29. The method of claim 23, wherein the neurological disease is selected from the group consisting of chronic dementia disease and Alzheimer's disease.
- 30. A method for prophylaxis or treatment of a neurological disease in a patient in need thereof, comprising the step of
 - administering to the patient a substance that causes modulation of the concentration of at least one VGF protein
 - or VGFARP peptide in a quantity sufficient to prevent or treat the neurological disease.

 31. The method of claim 30, wherein the neurological
- disease is selected from the group consisting of chronic dementia disease and Alzheimer's disease.

 32. The method of claim 30, wherein the modulation is a
- 32. The method of claim 30, wherein the modulation is a reduction in concentration of the at least one VGF protein or VGFARP peptide.
- 33. A method of claim 30, wherein the modulation is an increase in concentration of the at least one VGF protein or VGFARP peptide.
- 34. The method of claim 30, wherein the substance is selected from the group consisting of:
 - a) antibodies directed against VGF proteins, VGFARP peptides, NGF, BNDF or NT-3;
 - h) antisense nucleic acids, triplex nucleic acids or rihozymes that reduce expression of VGF proteins, VGFARP peptides, NGF, BNDF or NT-3;
 - c) substances that inhibit processing of VGF proteins; and
- d) antagonists of VGFARP peptides or VGF proteins.
 35. The method of claim 30, wherein the substance is selected from the group consisting of:
 - a) VGF proteins, VGFARP peptides, NGF, BNDF or NT-3;
 - b) nucleic acids which code for VGF proteins, VGFARP peptides, NGF, BNDF or NT-3;
 - substances which promote the processing of VGF proteins, and
- d) agonists of the VGFARP peptides or of VGF proteins.

 36. The method of claim 30, wherein the substance modulates the expression of at least one VGF protein.
- 37. The method of claim 36 wherein the VGF protein is selected from the group consisting of NGF, BNDF and NT.2
- 38. The method of claim 30, wherein the substance selectively inhibits or stimulates the transcription or expression of at least one VGF protein.

- The method of claim 30, wherein the substance binds to at least one VGF protein or VGFARP peptide.
- 40. The method of claim 39 wherein the substance is selected from the group consisting of antibodies, antibody fragments, and affinity matrices.
- 41. The method of claim 30 wherein the substance is administered via an administration route selected from the group consisting of: the bloodstream, the gastrointestinal tract, the urogenital tract, the lymphatic system, the sub-arachnoid space, the lungs, and direct injection into tissue.
- 42. The method of claim 41, wherein the tissue is selected from the group consisting of muscle tissue, adipose tissue, and brain tissue.
- 43. The method of claim 30, wherein the substance has been pharmaceutically processed or chemically or biologically modified to cross the blood-brain barrier and/or the blood-CSF barrier.
- 44. A VGFARP peptide.
- 45. The VGFARP peptide of claim 44, wherein the VGFARP peptide is a derivative of a VGF protein.

 46. The VGFARP peptide of claim 45, wherein the
- 46. The VGFARP peptide of claim 45, wherein t VGFARP peptide is a derivative of a VGF allele.
- 47. The VGFARP peptide of claim 44, wherein the VGFARP peptide is a mutant VGFARP peptide that differs by a maximum of 2 amino acids from the corresponding unmutated VGFARP peptide.
- 48. The VGFARP poptide of claim 44, wherein the sequence of the VGFARP poptide is selected from the group consisting of: SEQ ID NO.1, SEQ ID NO.2, SEQ ID NO.2, SEQ ID NO.3, SEQ ID NO.14, SEQ ID NO.11, SEQ ID NO.11, SEQ ID NO.11, SEQ ID NO.12, SEQ ID NO.13, SEQ ID NO.13, SEQ ID NO.13, SEQ ID NO.13, SEQ ID NO.14, SEQ ID NO.15, SEQ ID NO.15, SEQ ID NO.15, SEQ ID NO.15, SEQ ID NO.22, SEQ ID NO.23, SEQ ID NO.24, SEQ ID NO.25, SEQ I

- SEQ ID NO:29, SEQ ID NO:30, SEQ ID NO:31, SEQ ID NO:35, SEQ ID NO:35, SEQ ID NO:36, SEQ ID NO:40, SEQ ID NO:41, and SEQ ID NO:40, SEQ ID NO:41, and SEQ ID NO:40, SEQ ID NO:41, and SEQ ID NO:42, SEQ ID NO:42, SEQ ID NO:44, and SEQ ID NO:44, SEQ ID NO:45, SEQ ID NO:46, SEQ ID NO:46, SEQ ID NO:47, SEQ ID NO:47, SEQ ID NO:47, SEQ ID NO:48, SEQ ID NO:49, SEQ ID NO:40, SEQ ID NO:41, SEQ ID NO:40, SEQ ID NO:41, SEQ ID NO:45, SEQ ID NO:45,
- 49. The VGFARP peptide of claim 45, wherein said VGFARP peptide is a derivative of a VGF protein represented by SEQ ID NO:43 or SEQ ID NO:44.
- The VGFARP peptide of claim 44, wherein said peptide is chemically or post-translationally modified.
- peptide is chemically or post-translationally modified.

 51. A nucleic acid molecule that encodes a VGFARP peptide.
- 52. A nucleic acid molecule that is the complement of a nucleic acid molecule that encodes a VGFARP peptide.
- 53. A pharmaceutical composition comprising,
- at least one VGF protein or VGFARP peptide, a nucleic acid encoding at least one VGF protein or VGFARP peptide, or a nucleic acid that is the complement of a nucleic acid encoding at least one VGF protein or
- A diagnostic reagent for the detection of neurological diseases, comprising.
 - antibodies to at least one VGF protein or VGFARP peptide, and
 - a suitable carrier.

VGFARP peptide.

- 55. The diagnostic reagent of claim 54, wherein the neurological disease is selected from the group consisting of a neurological disease, chronic dementia, and Alzheimer's disease.
- 56. Antibodies that bind to VGFARP peptides.
- Nucleic acids that are VGF-specific antisense nucleic acids, components of VGF-specific ribozymes, or VGFspecific triplex nucleic acids.

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